wherein the pathway contains a protein with an APB domain, comprising the step of disrupting or promoting interaction between said APB domain and its binding partner *in vivo*.

- 2. (Amended) A method for treating a patient having a [disease or disorder] cancer or neoplasm characterized by APB binding comprising the step of administering to said patient a therapeutically effective amount of an agent which decreases binding between an APB recognition region present in a first protein and an APB domain present in a second protein.
- 3. (Amended) The method of claim 2, wherein said first protein is a receptor tyrosine kinase, said second protein is Shc, and said agent decreases one or more [activities of said receptor tyrosine] kinase <u>functions</u>
- 4. (Amended) The method of claim 3, wherein said receptor tyrosine kinase is selected from the group consisting of EGF receptor, HER-2, and Trk [TrkA].
- 20. (Amended) The method of claim 4 [or claim 19,] wherein said receptor tyrosine kinase is HER-2, and said [disease or disorder] <u>cancer or neoplasm</u> is breast cancer.
- 21. (Amended) The method of claim 4 [or claim 19,] wherein said [disease] cancer or neoplasm is cancer.
- 22. (Amended) The method of claim 4 [or claim 19,] wherein said receptor tyrosine kinase is [EFG] EGFR, and said [disease or disorder] cancer or neoplasm is at least one selected from the group consisting of gliomas, head cancers, neck cancers, gastric cancers, lung cancers, ovarian cancers, colon cancers, and prostate cancers.
- 23. (Amended) The method of claim 4 [or claim 19,] wherein said receptor tyrosine kinase is HER-2, and said [disease or disorder] cancer or neoplasm is at least one selected from the group consisting of stomach adenocarcinomas, salivary gland adenocarcinomas, endometrial

